IN THE CLAIMS:

All claims currently pending and under construction in the referenced application are shown below. Claims 7, 19, 20, 22, 24, 29, 34 and 36 are amended herein. New claims 37-40 are added. This listing of claims will replace all prior versions and listings of claims in the application. Claims 1-6, 8, 9, 12-18 and 25 and 26 were previously cancelled. Applicants respectfully submit that no new matter has been added.

Listing of Claims:

1-6. (Cancelled).

- 7. (Currently amended) An immunogenic composition for <u>allowing the marking of an exposure of a subject to wild-type Salmonella</u>, said immunogenic composition comprising:
- an immunologically effective amount of a live live mutated bacterium bacteria and a pharmaceutically acceptable carrier;
- wherein, said live mutated bacterium is bacteria are Salmonella enterica that in its their wild-type form carries-carried flagella having at least one antigenic determinant; and
- wherein <u>after mutation</u>, said live mutated <u>bacterium is bacteria are</u> not capable of inducing an immune response to <u>the</u> at least one antigenic determinant of flagellin <u>so as to allow marking of an exposure of the subject to the wild-type Salmonella in the subject to which it is administered.</u>
 - 8-9. (Cancelled).

- 10. (Previously Presented) The immunogenic composition according to claim 7, further comprising: an adjuvant selected from the group consisting of Freunds Complete adjuvant, Freunds Incomplete adjuvant, vitamin E, non-ionic block polymers, muramyldipeptides, immune stimulating complexes, saponins, mineral oil, vegetable oil, Carbopol, *E. coli* heat-labile toxin, *Cholera* toxin, aluminum hydroxide, aluminum phosphate, aluminum oxide, oil-emulsions, and vitamin-E solubilisate.
- 11. (Previously Presented) The immunogenic composition according to claim 7, wherein the immunogenic composition is in a freeze-dried or spray-dried form.
 - 12-18. (Cancelled).
- 19. (Currently amended) An immunogenic composition for <u>allowing the marking of</u> an exposure of a subject to wild-type *Salmonella*, said immunogenic composition comprising: an immunologically effective amount of an of inactivated mutated <u>bacterium bacteria</u>, the <u>inactivated mutated bacteria having a mutation in a gene encoding flagellin, and a pharmaceutically acceptable carrier;</u>
- wherein said inactivated mutated bacterium is bacteria are Salmonella enterica that in itstheir wild-type form earries-carried flagella having at least one antigenic determinant; and
- wherein said inactivated mutated bacterium is bacteria are not capable of inducing an immune response to the at least one antigenic determinant of flagellin in the a subject to which it is administered.

- 20. (Currently amended) A composition comprising:
- an immunologically effective amount of a of live mutated Salmonella typhimurium, wherein the wild-type form of the live mutated S. typhimurium earries carried flagella having at least one antigenic determinant;
- wherein said live mutated S. typhimurium is are not capable of inducing an immune response to the at least one antigenic determinant of flagellin in a subject to which it is administered; and
- a pharmaceutically acceptable carrier comprising water, a solution of physiological salt concentration, SPGA, sorbitol, mannitol, starch, sucrose, glucose, dextran, albumin, casein, bovine serum, skim milk, or phosphate buffer.
 - 21. (Previously Presented) A composition comprising:
- an immunologically effective amount of a of mutated Salmonella typhimurium, wherein the wild type form of the mutated S. typhimurium carries flagella;
- wherein said mutated S. typhimurium is lacking lack flagellin and comprises comprise an immunologically effective amount of a S. typhimurium strain STMP mutated bacterium; and
- a pharmaceutically acceptable carrier.
- 22. (Currently amended) An improved Salmonella vaccine, having an immunologically effective amount of a of Salmonella enterica bacterium bacteria and an adjuvant in a pharmaceutically acceptable carrier, the improvement comprising:
- the Salmonella enterica bacterium bacteria comprising an inactivated mutated bacterium that in its their wild type form earries carried flagella having at least one antigenic determinant, but in its their mutated form is no longer capable of inducing an immune response to the at least one antigenic determinant of flagellin in a subject to which it the vaccine is administered, the mutated form having a mutation in a gene encoding flagellin.
 - 23. (Cancelled).

- 24. (Currently amended) The improved Salmonella vaccine of claim 22, wherein the inactivated mutated bacterium bacteria lacks flagellin.
 - 25-27. (Cancelled).
- 28. (Previously Presented) The improved *Salmonella* vaccine of claim 22, wherein the improved *Salmonella* vaccine is in a freeze-dried or spray-dried form.
- 29. (Currently amended) An improvement in a A marker vaccine, comprising a comprising Salmonella enterica bacterium bacteria, the improvement marker vaccine comprising:
- an immunologically effective amount of a of mutated Salmonella enterica, wherein the wild type form of the mutated Salmonella enterica carries carried flagella having at least one antigenic determinant;
- wherein said mutated Salmonella enterica bacterium is bacteria have a mutation in a gene encoding flagellin and are not capable of inducing an immune response to the at least one antigenic determinant of flagellin in a subject to which it is administered;

an adjuvant;

a pharmaceutically acceptable carrier; and wherein the marker vaccine is in a freeze-dried or spray-dried form.

- 30. (Previously Presented) The immunogenic composition according to claim 19, wherein the immunogenic composition is in a freeze-dried or spray-dried form.
- 31. (Previously Presented) The improved marker vaccine of claim 29, wherein the mutated *Salmonella enterica* is in live attenuated form.
- 32. (Previously Presented) The improved marker vaccine of claim 29, wherein the mutated Salmonella enterica lacks flagellin.

- 33. (Cancelled).
- 34. (Currently amended) In an immunogenic composition including a *Salmonella* bacterium, the improvement comprising:
- a lyophilized immunogenic composition comprising a mutated Salmonella enterica;
- said Salmonella enterica in its wild type form carrying flagella having at least one antigenic determinant; and
- said mutated Salmonella enterica lacking at least one antigenic determinant of flagellin and not being capable of inducing an immune response to the at least one antigenic determinant of flagellin in a subject to which it is administered, the mutated Salmonella enterica having a mutation in a gene encoding flagellin.
 - 35. (Cancelled).
 - 36. (Currently amended) A composition comprising:
- an immunologically effective amount of a of mutated S. typhimurium, wherein the wild type form of the mutated S. typhimurium carries flagella;
- wherein said mutated S. typhimurium comprises an immunologically effective amount of a of S. typhimurium strain STMP mutated bacterium bacteria; and a pharmaceutically acceptable carrier.
- 37. (New) An immunogenic composition for allowing the marking of an exposure of a subject to wild-type *Salmonella*, said immunogenic composition consisting essentially of: an immunologically effective amount of live mutated bacteria and water; an adjuvant;
- wherein, said live mutated bacteria are Salmonella enterica that in their wild-type form carried flagella having at least one antigenic determinant; and
- wherein after mutation said live mutated bacteria are not capable of inducing an immune response to the at least one antigenic determinant of flagellin so as to allow marking of an exposure of the subject to the wild-type Salmonella in the subject to which the vaccine is

administered.

- 38. (New) An immunogenic composition for allowing the marking of an exposure of a subject to wild-type *Salmonella*, said immunogenic composition consisting essentially of: an immunologically effective amount of live mutated bacteria and water;
- wherein, said live mutated bacteria are Salmonella enterica that in their wild-type form carried flagella having at least one antigenic determinant; and
- wherein after mutation said live mutated bacteria are not capable of inducing an immune response to the at least one antigenic determinant of flagellin so as to allow marking of an exposure of the subject to the wild-type Salmonella in the subject to which the immunogenic composition is administered.
- 39. (New) An immunogenic composition for allowing the marking of an exposure of a subject to wild-type *Salmonella*, said immunogenic composition comprising:
- an immunologically effective amount of live mutated bacteria and a pharmaceutically acceptable carrier;

an adjuvant;

- wherein, said live mutated bacteria are Salmonella enterica that in their wild-type form carried flagella having at least one antigenic determinant; and
- wherein after mutation said live mutated bacteria are not capable of inducing an immune response to the at least one antigenic determinant of flagellin so as to allow marking of an exposure of the subject to the wild-type *Salmonella* in the subject to which the immunogenic composition is administered.
 - 40. (New) A composition consisting essentially of:
- an immunologically effective amount of live mutated *Salmonella typhimurium*, wherein the wild-type form of the live mutated *S. typhimurium* carried flagella having at least one antigenic determinant;
- wherein said live mutated S. typhimurium are not capable of inducing an immune response to the at least one antigenic determinant of flagellin in a subject to which the immunogenic

composition is administered; and

a pharmaceutically acceptable carrier comprising water, a solution of physiological salt concentration, SPGA, sorbitol, mannitol, starch, sucrose, dextran, albumin, casein, bovine serum, skim milk, or phosphate buffer.